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(71) Applicant (for all designated States except US): TI-  
BOTEC PHARMACEUTICALS LTD. [IE/IE]; Little  
Island, Co Cork (IE).

(72) Inventors; and

(75) Inventors/Applicants (for US only): SIMMEN, Ken-  
neth Alan [GB/BE]; Boterbloemenlaan 35, B-3080

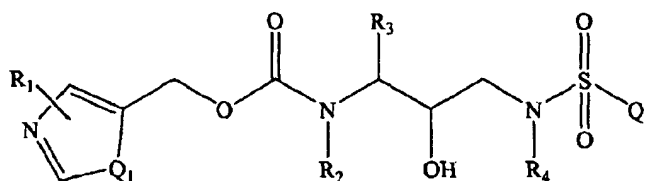
Tervuren (BE). VAN ACKER, Koenraad Lodewijk  
August [BE/BE]; Uilenberg 30, B-9140 Temse (BE).  
WIGERINCK, Piet Tom Bert Paul [BE/BE]; Kardinaal  
Cardijnstraat 29, B-2840 Terhagen (BE). SURLERAUX,  
Dominique Louis Nestor Ghislain [BE/BE]; rue de la  
scaillée, 13, B-1440 Braine-le-château (BE). DAMS,  
Géry Karel Julia [BE/BE]; Nieuwe Erven 11, B-3583  
PAAL-BERINGEN (BE). QUIRYNEN, Ludo Maria  
Marcel [BE/BE]; Katelijnestraat 19F, B-2320 Hoogstraten  
(BE). HERTOOGS, Kurt [BE/BE]; Albert I laan 185  
bus 401, B-8260 Nieuwpoort (BE). PAUWELS, Rudi  
Wilfried Jan [BE/CH]; Chemin de Layaz 3, CH-1806  
Saint-Légier La Chiésaz (CH).

(74) Agent: DAELEMANS, Frank; Tibotec-Virco Comm.  
VA, Generaal De Wittelaan L 11B 3, B-2800 Mechelen  
(BE).

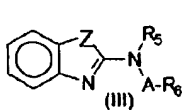
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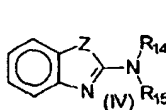
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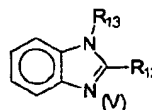
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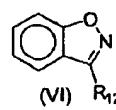
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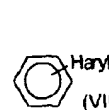
(IV)



(V)



(VI)



(VII)

(57) Abstract: The present invention concerns sulfonamide derivatives having the general formula (I) and *N*-oxides, salts, stereoisomeric forms, racemic mixtures, prodrugs and esters thereof, wherein Q<sub>1</sub> is -S- or -O-; R<sub>1</sub> is hydrogen, C<sub>1-6</sub>alkyl, hydroxy, amino, halogen, aminoC<sub>1-4</sub>alkyl and mono- or di(C<sub>1-4</sub>alkyl)amino; R<sub>2</sub> is hydrogen or C<sub>1-6</sub>alkyl; R<sub>3</sub> is C<sub>1-6</sub>alkyl, aryl, C<sub>3-7</sub>cycloalkyl, C<sub>3-7</sub>cycloalkylC<sub>1-4</sub>alkyl, or arylC<sub>1-4</sub>alkyl; R<sub>4</sub> is hydrogen, C<sub>1-4</sub>alkyloxycarbonyl, carboxyl, optionally mono- or disubstituted aminocarbonyl, mono- or di(C<sub>1-4</sub>alkyl)aminocarbonyl, C<sub>3-7</sub>cycloalkyl, C<sub>2-6</sub>alkenyl, C<sub>2-6</sub>alkynyl or C<sub>1-6</sub>alkyl optionally substituted with one or more substituents each independently selected from aryl, Het<sup>1</sup>, Het<sup>2</sup>, C<sub>3-7</sub>cycloalkyl, C<sub>1-4</sub>alkyloxy-carbonyl, carboxyl, aminocarbonyl, mono- or di(C<sub>1-4</sub>alkyl)aminocarbonyl, aminosulfonyl, C<sub>1-4</sub>alkylS(=O)<sub>n</sub>, hydroxy, cyano, halogen or amino optionally mono- or di-substituted where the substituents are each independently selected from C<sub>1-4</sub>alkyl, aryl, arylC<sub>1-4</sub>alkyl, C<sub>3-7</sub>cycloalkyl, C<sub>3-7</sub>cycloalkylC<sub>1-4</sub>alkyl, Het<sup>1</sup>, Het<sup>2</sup>, Het<sup>1</sup>C<sub>1-4</sub>alkyl and Het<sup>2</sup>C<sub>1-4</sub>alkyl; Q<sub>2</sub> is a radical of formulae (III), IV, V, VI, VII) for the manufacture of a medicament useful for inhibiting HCV activity in a mammal infected with HCV. The present invention also relates to the use of said sulfonamides in pharmaceutical compositions aimed to treat or combat combined HCV and HIV infections. In addition, the present invention relates to processes for preparation of such pharmaceutical compositions. The present invention also concerns combinations of the present sulfonamides with other anti-HCV agents and/or anti-HIV agents.

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